### Amendments to the SPECIFICATION

## TITLE OF INVENTION

Dengue virus mutant, strain MBU 01-2002

# DENGUE VIRUS MUTANT, STRAIN MBU 01-2002

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] Not Applicable This application claim priority to Thai Application No. 075425 filed July 26, 2002.

STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH OR

DEVELOPMENT

—— Not Applicable

REFERENCE TO SEQUENCE LISTING, A TABLE, OR A COMPUTER PROGRAM LISTING COMPACT DISC APPENDIX

— Not Applicable

### FIELD OF THE INVENTION

[0002] The present invention relates to dengue virus containing a full genome-length dengue virus genome with a defined mutation.

### BACKGROUND OF THE INVENTION

[0003] Dengue virus is a causative agent of dengue fever and dengue hemorrhagic fever, which are widespread in several tropical countries around the world. Transmission of dengue virus among humans occurred through mosquito biting activity; infected human serve principally as the reservoirs for transmission in the urban setting. Among viral isolates obtained from infected humans and mosquitoes, four serologically defined types (serotypes) of dengue virus are known, all of which can cause potentially fatal dengue hemorrhagic fever (D.S. Burke and T.P. Monath (2001) Flaviviruses, P. 1043-1125, in D.M. Knipe, P.M. Howley, D.E. Griffin, R.A. Lamb, M.A. Martin, B. Roizman and S.E. Strauss (ed.), Fields Virology, 4<sup>th</sup> ed. Lippincott Williams & Wilkins, Philadelphia, PA). In human, primary infection by dengue virus of any serotype

generally causes milder diseases and induces immunity only against the infecting serotype. As cross protective immunity is short-lived, secondary infection by other dengue serotypes is common in countries where two or more serotypes co-circulate. It is well accepted that secondary infection increases the risk of developing dengue hemorrhagic fever. Effective prevention of dengue hemorrhagic fever will require a vaccine, which can induce protective immunity against all four serotypes of dengue virus.

[0004] Dengue virus is a member of the Genus Flavivirus in the Family Flaviviridae. The virion is spherical in shape with the diameter of about 50 nm. The outer part of the virion (envelope) consists of lipid bilayer and two different glycoproteins, E and prM/M (B.D. Lindenbach and C.M. Rice. (2001) Flaviviridae: the viruses and their replication, p. 991-1041 in D.M. Knipe, P.M. Howley, D.E. Griffin, R.A. Lamb, M.A. Martin, B. Roizman and S.E. Strauss (ed.), Fields Virology, 4<sup>th</sup> ed. Lippincott Williams & Wilkins, Philadelphia, PA). The core is composed of another protein, C, in association with the single-stranded RNA genome. In each virion, one molecule of about 10.7 kb long RNA genome is present; it encodes three structural proteins and seven non-structural proteins, which are required for virus multiplication inside the infected cells, but are not components of virion. The organization of the genome is as follows: 5'cap-5'untranslated region-C-prM/M-E-NS- NS2A-NS2B-NS3-NS4A-NS4B-NS5-3' untranslated region-3'end. This genome organization is common to all flaviviruses.

[0005] Following the assembly of virion in the endoplasmic reticulum, the envelope of immature virions, consisting of only prM and E proteins, is further modified by glycosylating enzymes in the Golgi apparatus and by the proteolytic enzyme, furin, in the *trans*-Golgi network. Specifically, when immature virion is exported along the secretory pathway to the extracellular milieu, furin cleaves the protein prM internally at the pr-M junction, generating virion-associated M protein and soluble pr peptide, which no longer associate with the virion (K. Stadler, et al. (1997) Proteolytic activation of tick-borne encephalitis virus by furin. J. Virol. 71:8475-8481). Cleavage of prM is absolutely necessary for the ability of the mature, extracellular virion of flavivirus to initiate productive infection of the host cell (S. Elshuber, et al. (2003) Cleavage of protein prM is necessary for infection of BHK-21 cells by tick-borne encephalitis virus. J. Gen. Virol. 84:183-191). However, cleavage of prM can be incomplete and

extracellular virions of several flaviviruses were known to contain varying amount of prM protein. The significance of the remaining prM on the envelope is not yet known.

[0006] Recent structural data of the immature particles of dengue virus reveal that the prM proteins associate with the E proteins as sets of prM-E heterodimer (Y. Zhang, et al. (2003) Structures of immature flavivirus particles.—EMBO J. 22:2604-13). The prM proteins project out of the surface whereas the E proteins lie flat and parallel to the lipid bilayer of the envelope. The protruding portion of the prM proteins covers the tip of the E protein that is responsible for fusing activity of E protein. The structure of a preparation of the mature dengue virion in which all prM proteins are cut is quite different from the immature virion including the formation of E-E homodimers (R.J. JKuhn, et al. (2002) Structure of dengue virus: implications for flavivirus organization, maturation, and fusion. Cell, 108:717-25), indicating that cleavage of prM by furin leads to a significant rearrangement of the envelope proteins. Such rearrangement must be important for generating infectious viral particles.

[0007] It has been repeatedly observed that the extracellular virions of dengue virus usually contain remaining prM protein on their envelope (R. Anderson, et al. (1997) Activation of endothelial cells via antibody-enhanced dengue virus infection of peripheral blood monocytes. J. Virol. 71: 4226-4232; R.T. He, et al. (1995) Antibodies that block virus attachment to Vero cells are a major component of the human neutralizing antibody response against dengue virus type 2. J. Med. Virol. 45: 451-461; E.A. Henchal et al. (1985) Epitope analysis of antigenic determinants on the surface of dengue-2 virions using monoclonal antibodies. Am. J. Trop. Med. Hyg. 34: 162-169; J.M. Murray et al. (1993) Processing of the dengue virus type 2 proteins prM and CprM. J. Gen. Virol. 74: 175-182; V.B. Randolph et al. (1990) Acidetropic amines inhibit proteolytic processing of flavivirus prM-protein. Virology 174: 450-458; J.T. Roehrig et al. (1998) Monoclonal antibody mapping of the envelope glycoprotein of the dengue 2 virus, Jamaica. Virology 246: 317-328; S. Wang et al. (1999) PrM-and cell-binding domains of the dengue virus E protein. J. Virol. 73: 2547-2551). The observations were made in dengue virus prepared from a number of cell lines of both mosquito and mammalian origins in several laboratories. These dengue virus preparations are infectious and able to generate new rounds of infection efficiently.

[0008] When compared with other flaviviruses in which the prM protein is completely cleaved, an incomplete cleavage of prM in dengue virus coincides with a lower number of positively charged amino acids and also the presence of two negatively charged amino acids within the 13-amino acid, pr-M junction proximal sequence, which extends beyond the P6 and P4 boundary previously known to affect cleavage of target protein by furin (G. Thomas (2002) Furin at the cutting edge: from protein traffic to embryogenesis and disease. Nat. Rev. Mol. Cell. Biol. 3: 753-766; K. Nakayama (1997) Furin: a mammalian subtilisin/Kex2p-like endoprotease involved in processing of a wide variety of precursor proteins. Biochem. J. 327: 625-635; A. Zhou et al. (1999) Proteolytic processing in the secretory pathway. J. Biol. Chem. 274: 20745-20748). Thus, one of the distinctive features of dengue virus is the conservation of the pr-M cleavage junction sequence, which allows only partial cleavage by host cell-derived furin. Alterations of dengue pr-M cleavage junction may modify the structural characteristics of the virion and the biology of dengue virus, especially the replication kinetics in ways that are not yet known. Alterations of this pr-M junction sequence in dengue virus can take a number of ways, including: a substitution of each amino acid position to alter one at a time the charge characteristics and the size of the R group; a substitution of two- or more amino acid positions in various combinations, a deletion of one or more charged amino acid positions; an insertion of one or more charged amino acids or any combination of these manipulation methods. Depending on the nature of the amino acid changes and the property of furin in specific cell lines tested, the cleavage of prM protein in the mutant dengue viruses can either be enhanced, lowered, or unaffected. Alteration of the cleavage of the N-terminal of prM protein of a related flavivirus, Yellow fever virus, has been known to reduce virus replication by affecting the production of virus within the infected cells (E. Lee et al. (2000) Mutagenesis of the signal sequence of yellow fever virus prM protein: enhancement of signalase cleavage in vitro is lethal for virus production. J. Virol. 74: 24–32).

C. Lai et al.; Proceedings of the National Academy of Sciences USA, vol. 88, Jun. 1991;

Infectious RNA transcribed from stably cloned full length cDNA of dengue type 4 virus; pp. 5139-5143.

S. Elshuber et al.; Journal of General Virology, vol. 84, 2003; Cleavage of protein prM is necessary for infection of BHK-21 cells by tick-borne encephalitis virus; pp. 183-191.

R. Sriburi et al.; Journal of Virological Methods, vol. 92, 2001; Construction of infectious dengue 2 virus cDNA clones using high copy number plasmid; pp. 71-82.

R.M. Kinney et al.; Virology, vol. 230, 1997; Construction of infectious cDNA clones for dengue 2 virus: strain 16681 and its attenuated vaccine derivative, strain PDK-53; pp. 300-308.

E. Lee et al.; Journal of Virology, vol. 74, 2000; Mutagenesis of the signal sequence of yellow fever virus prM protein; enhancement of signalase cleavage *in vitro* is lethal for virus production; pp. 24-32.

K. Stadler et al.; Journal of Virology, vol. 71, 1997; Proteolytic activation of tick borne encephalitis virus by furin; pp. 8475–8481.

Y. Zhang et al.; European Molecular Biology Organization Journal, vol. 22, 2003; Structure of immature flavivirus particles; pp. 2604-2613.

R.J. Kuhn et al.; Cell, vol. 108, 2002; Structure of dengue virus: implications for flavivirus organization, maturation, and fusion; pp. 717-725.

R. Anderson et al.; Journal of Virology, vol. 71, 1997; Activation of endothelial cells via antibody enhanced dengue virus infection of peripheral blood monocytes; pp. 4226-4232.

R.T. He et al,; Journal of Medical Virology, vol. 45; 1995, Antibodies that block virus attachment to Vero cells are a major component of the human neutralizing antibody response against dengue virus type 2; pp. 451-461.

E.A. Henchal et al.; American Journal of Tropical Medicine and Hygiene; vol. 34, 1985; Epitope analysis of antigenic determinants on the surface of dengue 2 virions using monoclonal antibodies; pp. 162-169.

J.M. Murray et al.; Journal of General Virology, vol. 74, 1993. Processing of the dengue virus type 2 proteins prM and C prM; pp. 175-182.

V.B. Randolph et al.; Virology, vol. 174, 1990; Acidotropic amines inhibit proteolytic processing of flavivirus prM protein; pp. 450-458.

J.T. Roehrig et al.; Journal of Virology, vol. 73, 1998; Monoclonal antibody mapping of the envelope glycoprotein of the dengue 2 virus, Jamaica; pp. 317-328.

S. Wang et al.; Journal of Virology, vol. 73, 1999; PrM- and cell-binding domains of the dengue virus E protein; pp. 2547-2551.

G. Thomas; Nature Review of Molecular Cell Biology, vol. 3, 2002; Furin at the cutting edge: from protein traffic to embryogenesis and disease; pp. 753–766.

K. Nakayama; Biochemical Journal, vol. 327, 1997; Furin: a mammalian subtilisin/Kex2p-like endoprotease involved in processing of a wide variety of precursor proteins; pp. 625-635.

A. Zhou et al.; Journal of Biological Chemistry, vol. 274, 1999; Proteolytic processing in the secretory pathway; pp. 20745-20748.

### BRIEF SUMMARY OF THE INVENTION

[0009] The present invention is a genetic construct and a dengue virus containing a full genome-length dengue virus genome with a defined mutation within the 13-amino acid-coding region proximal to the pr-M cleavage junction. The mutation involves increasing the number of positively charged amino acid acids and abolishing the negatively charged amino acid acids of the pr-M cleavage junction, which results in an enhanced cleavage of the prM protein to the M protein. The mutant dengue virus possesses less prM protein on the viral envelope than the prototype dengue virus. The mutant dengue virus was able to bind, enter and replicate inside the infected cells as efficiently as the prototype dengue virus, but was ineffective in its release from infected cells into the extracellular compartment, causing a greatly reduced virus titer and

replication kinetics. The mutant dengue virus strain MBU01-2002 may be useful as an immunogen for the generation of protective immunity against dengue virus infection or dengue hemorrhagic fever in human in the future.

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWING
Not Applicable

# LISTING [00010] SEQ ID NO: 1 is the genomic nucleotide sequence of Dengue virus. [00011] SEQ ID NO: 2 is primer which can be used to introduce specific mutant sequence into the pr-M junction of dengue genome (sense primer: 5' TATGGACGGTGCACGCGGACCAGGCATTCCAAGAGATCTAGGA 3'). [00012] SEQ ID NO: 3 is primer which can be used to introduce specific mutant sequence into the pr-M junction of dengue genome (anti-sense primer: 5' GATCTCCTAGATCTCTTGGAATGCCTGGTCCGCGTGCTCCA 3').

# DETAILED DESCRPTION OF THE INVENTION

1. Generation of a mutant dengue virus strain MBU 01-2002

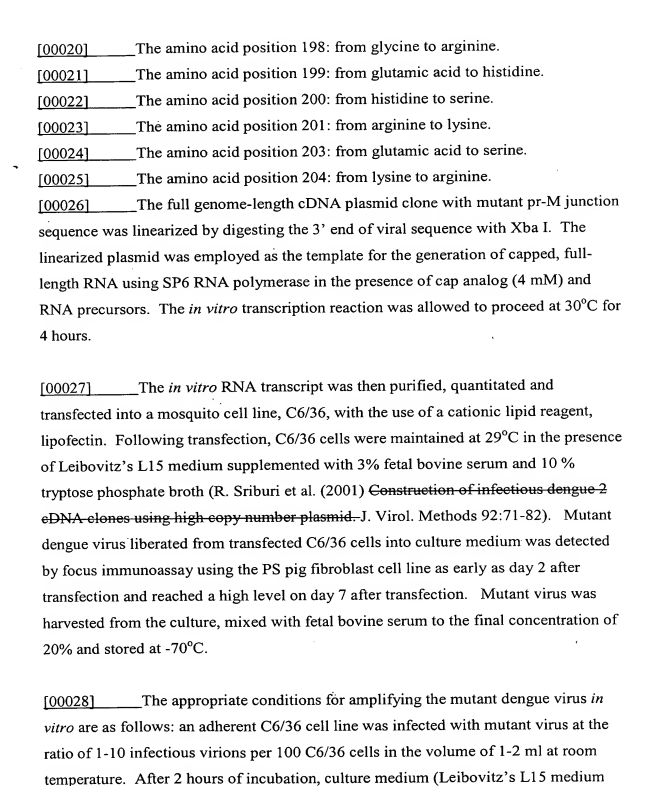
[00013] \_\_\_\_\_ Dengue virus mutant strain MBU 01-2002 was generated by altering a specific gene segment of a virulent, prototype dengue virus, dengue serotype 2 virus strain 16681, a wild type strain, also called prototype dengue virus. which The virus strain 16681 was isolated from a Thai patient with dengue hemorrhagic fever. A change was introduced specifically at the pr-M junction, just proximal to the cleavage site of a host proteolytic enzyme (furin), within the coding region for the prM protein. The alteration was performed in successive steps, as follows:

A plasmid subclone containing the sequence nt 1- 1547 of a full-length cDNA clone of dengue virus strain 16681 (R. Sriburi et al. (2001) Construction of infectious dengue 2 virus cDNA clones using high copy number plasmid. J. Virol. Methods 92:71-82) was altered by introducing substitution mutation at two nucleotide positions, 666 (T666A mutation) and 709 (A709G mutation), using a PCR-based, site-directed mutagenesis scheme. The alteration resulted in new restriction enzyme recognition sites for Nde I and BamH I at the nucleotide positions 666 and 709 of the

viral sequence, respectively. The presence of the new recognition sites was confirmed by nucleotide sequence analysis of the resultant plasmid subclones. \_In order to introduce specific mutant sequence into the pr-M junction of dengue genome, two single-stranded oligonucleotides, SEQ ID NO: 2 (sense primer: 5' TATGGACGGTGCACGCGGACCAGGCATTCCAAGAGATCTAGGA 3') and SEQ ID NO: 3 (anti-sense primer: 5' GATCTCCTAGATCTCTTGGAATGCCTGGTCCGCGTGCTCCGTCCA 3'), containing the following sequences were designed and synthesized. 5' TATGGACGGTGCACGCGGACCAGGCATTCCAAGAGATCTAGGA 3' (sense <del>primer)</del> 5' GATCTCCTAGATCTCTTGGAATGCCTGGTCCGCGTGCTCCA 3' (antisense primer) The oligonucleotides were mixed and allowed to anneal at 37°C to generate a short double-stranded oligonucleotide with a Nde I-compatible end and a BamH I-compatible end. The annealed, double-stranded oligonucleotide was then ligated into the plasmid subclone mentioned above, which was previously digested with Nde I and BamH I enzyme and was devoid of a short Nde I-BamH I fragment. Following transformation of the ligated product into E. coli and the selection with ampicillin, the presence of mutant sequence in the resultant subclones was confirmed by nucleotide sequence analysis. The mutant pr-M junction sequence was then introduced into the plasmids containing the 5' half-genome (nt positions 1-4497) and subsequently the full-length cDNA clone (nt positions 1-10723). The final full-length cDNA plasmid contained the intended mutant sequence at the pr-M junction, which encodes a mutant prM protein with the following amino acid changes acids change (numbering is according to the sequence of the polyprotein of dengue virus serotype 2 as described in R. M. Kinney et al. (1997) Construction of infectious cDNA clones for dengue 2 virus: strain 16681 and its attenuated vaccine derivative, strain PDK-53. Virology 230: 300-308): The amino acid position 193: from threonine to arginine. [00017] The amino acid position 196: from threonine to arginine. [00018]

The amino acid position 197: from methionine to threonine.

[00019]



supplemented with 3% fetal bovine serum and 10 % tryptose phosphate broth) was added to 15 ml and the cell line was maintained at 29°C and the liberated virus was

monitored daily. When high virus level was reached, the mutant virus was harvested, mixed with fetal bovine serum to the concentration of 20% and stored in small aliquots

at -70°C.

1.

2. Properties of the mutant dengue virus strain MBU 01-2002
[00029] The mutant dengue virus strain MBU 01-2002 is 10-1,000 folds less
efficient in the replication in vitro than the prototype dengue virus strain 16681 when
tested in the C6/36 mosquito cell line, PS pig fibroblast cell line, Vero monkey kidney
cell line and human embryonic kidney 293T cell line.
[00030] The extracellular virions of the mutant dengue virus strain MBU 01-2002
contain less prM protein on the viral envelope than the prototype dengue virus strain
16681 when tested by immunoblot analysis of unlabelled virions, and by SDS-PAGE
analysis of specifically immunoprecipitated, S35-labelled virions. Reduced level of prM
on virion envelope indicates higher efficiency of the cleavage of pr-M junction during
virus export through the trans-Golgi network of infected cells. By design, the remaining
prM protein on the envelope of the mutant dengue virus strain MBU01-2002 contains a
higher number of positively-charged amino acids than those of the prototype dengue
virus strain 16681.
[00031] The mutant dengue virus strain MBU 01-2002 is capable of inducing
infected C6/36 mosquito cell line to fuse at both neutral condition (pH=7.0) and acidic
condition (pH less than 7.0) whereas the parent 16681 dengue virus requires acidic
condition. Induction of infected cell fusion by the mutant dengue virus strain MBU 01-
2002 occurs well at 29°C, but is less efficient at 40°C.
The mutant dengue virus strain MBU 01-2002 binds and enters PS cell
line and 293T cell line as efficiently as the prototype dengue virus strain 16681. Within
the infected cells, the mutant dengue virus strain MBU 01-2002 replicates as efficiently
as the prototype dengue virus strain 16681, but was exported out of the infected cells to
a lesser extent than the prototype dengue virus.
[00033] The invention provides a significant and unexpected improvement to the
art of mutant dengue virus production. One with skill in the art could expect that a
mutant dengue virus with enhanced cleavage of the prM protein to be more (or, at least
as) efficient in the replication in cultured cell lines as the prototype virus. However,
unexpectedly, the present invention identifies a mutant dengue virus with defective

release out of the infected cells, causing the lower virus titer and lower growth kinetics in cultured cell lines. The mutant dengue virus also differs from the prototype dengue virus in its ability to induce infected mosquito cell fusion at neutral pH upon incubation at 29°C. Because of its lowered replication kinetics, the invention has a potential to serve as a live, attenuated vaccine candidate for inducing protective immune response against dengue virus in humans.